



Microgeographic adaptation and the effect of pollen flow on the adaptive potential of a temperate tree species

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Summary

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• In species with long-distance dispersal capacities and inhabiting a large ecological niche, local selection and gene flow are expected to be major evolutionary forces affecting the genetic adaptation of natural populations. Yet, in species such as trees, evidence of microgeographic adaptation and the quantitative assessment of the impact of gene flow on adaptive genetic variation are still limited.

• Here, we used extensive genetic and phenotypic data from European beech seedlings collected along an elevation gradient, and grown in a common garden, to study the signature of selection on the divergence of eleven potentially adaptive traits, and to assess the role of gene flow in resupplying adaptive genetic variation.

• We found a significant signal of adaptive differentiation among plots separated by < 1 km, with selection acting on growth and phenological traits. Consistent with theoretical expectations, our results suggest that pollen dispersal contributes to increase genetic diversity for these locally differentiated traits.

• Our results thus highlight that local selection is an important evolutionary force in natural tree populations and suggest that management interventions to facilitate movement of gametes along short ecological gradients would boost genetic diversity of individual tree populations, and enhance their adaptive potential to rapidly changing environments.

Introduction

Recent interest in microgeographic adaptation - adaptation at spatial scales compatible with substantial amounts of gene dispersal - suggests that we must reconsider the scale at which evolution occurs (Richardson et al., 2014). At this fine spatial scale, a major question that remains to be answered is whether gene flow is constraining or facilitating local adaptation. Too much gene flow would overwhelm the effects of natural selection that drive adaptive genetic divergence among populations (Lenormand, 2002) and decrease the adaptation of populations to their local environment (migration load). Conversely, gene flow also can resupply the genetic diversity eroded by genetic drift and selection, and thus facilitate future evolutionary responses (Pease et al., 1989; Barton, 2001; Lenormand, 2002; Bridle et al., 2010; Polechova & Barton, 2015). In many taxa, and especially long-lived organisms, gene flow is expected to be the main process maintaining the standing genetic variation within natural populations (Barton, 2001; Le Corre & Kremer, 2003; Bridle et al., 2010). In the particular case of environments varying through both space and time, gene flow also could spread alleles pre-adapted to future environmental conditions (Davis & Shaw, 2001; Aitken et al., 2008; Kremer et al., 2012; Fitzpatrick & Reid, 2019). Although theoretical developments on this topic have been rich and

© 2020 The Authors New Phytologist © 2020 New Phytologist Trust provided interesting hypotheses to test in nature, empirical studies on the role and importance of gene flow in adaptation are still lacking (but see Fitzpatrick *et al.*, 2015, 2016).

Increasing evidence of microgeographic adaptation in nature provides strong support for the idea that gene flow rarely prevents the emergence of local adaptation or adaptive divergence (Anderson et al., 2015; Eckert et al., 2015; Fitzpatrick et al., 2015; Moody et al., 2015; Peterson et al., 2016; for a review, see Richardson et al., 2014). This evidence strengthens the conclusions of studies at larger spatial scales, showing that geographic or genetic distances among populations are not correlated with the level of local adaptation or adaptive genetic divergence (McKay & Latta, 2002; Leimu & Fischer, 2008; Hereford, 2009). These results suggest that selective pressures can be strong enough to counterbalance the effects of gene flow, and to shape genetic variation at a fine spatial scale. However, evidence of microgeographic adaptation is reported mainly for certain taxa (e.g. fish, Fitzpatrick et al., 2015; Moody et al., 2015; and annual or perennial herbs, Anderson et al., 2015; Peterson et al., 2016), whereas others, such as temperate tree species, are under-represented. In trees, there is a long tradition of large-scale adaptation studies, through provenance trials, following a quantitative genetic approach (Savolainen et al., 2007; Alberto et al., 2013). The development of genome-wide data and statistical methods has

considerably changed this research field, and evidence of adaptations at regional and landscape scales are now flourishing (e.g. Csillery *et al.*, 2014; Pluess *et al.*, 2016). Yet, the adaptive response of phenotypic traits at the local scale is still poorly understood (but see Brousseau *et al.*, 2013; Eckert *et al.*, 2015). Methodological limitations that previously hampered the robust testing of local phenotypic adaptation have been largely overcome (Ovaskainen *et al.*, 2011), permitting more accurate assessments of the scale at which phenotypic adaptations occur.

Empirical evidence of beneficial effects of gene flow on adaptive potential in nature are still rare, and often are assessed in laboratory settings (e.g. Swindell & Bouzat, 2006). Some experimental studies on inbred plant populations have demonstrated the beneficial effect of gene flow on the reduction of homozygosity and reintroduction of variation for fixed deleterious alleles (e.g. Costa e Silva et al. 2014; Bontrager & Angert, 2019). However, for outbred species, the role of gene flow on the maintenance of genetic diversity has been difficult to demonstrate, especially when relying on natural crosses. This effect of gene flow is thought to be particularly important when the populations exchanging genes are strongly genetically differentiated (Pease et al., 1989; Barton, 2001). Long-distance dispersal events could therefore play a major role in resupplying the genetic variation of populations and favour the spread of beneficial alleles (Savolainen et al., 2007; Kremer et al., 2012). In wind-pollinated species (anemophilous), these long-distance dispersal events are common, with mean pollen dispersal distances that can easily reach hundreds of metres, with rare events of spread over tens of kilometres in some species (Austerlitz et al., 2004; Petit & Hampe, 2006; Ashley, 2010; Kremer et al., 2012). Pollen dispersal may thus be highly efficient for spreading beneficial alleles between populations (Kremer *et al.*, 2012). Even though a pollen grain brings half as many immigrating alleles as does a seed, gene flow by pollen or seed is expected to have similar consequences on a population's adaptation and migration load (Lopez *et al.*, 2008).

Wind-pollinated tree species are therefore good study systems to look for empirical evidence of the beneficial effects of long-distance gene flow on adaptive capacities in nature. However, to our knowledge, only two studies have looked for such evidence. In Pinus contorta, Yeaman & Jarvis (2006) showed a positive correlation between the level of genetic variance within populations for growth and the spatial heterogeneity of selection. This pattern is consistent with the theoretical expectation that gene flow among differentiated populations is an important source of genetic variation. Along a latitudinal gradient of genetically differentiated populations of Pinus sylvestris, Nilsson (1995) found that offspring harvested after one episode of reproduction showed a shift in average genetic values as compared to their mothers. This pattern is expected in the case of directional gene flow among locally differentiated populations (Fig. 1). This effect of directional pollen dispersal on an offspring's average genetic values for adaptive traits could be particularly beneficial if the environmental conditions of the recipient population change toward the one of the donor population (Davis & Shaw, 2001; Aitken et al., 2008).

The aim of the present study is to investigate the effects of natural selection and long-distance gene flow on the adaptive potential of a major temperate tree species, the European beech (*Fagus sylvatica* L.). This anemophilous species can disperse pollen at long distance (Piotti *et al.*, 2011), a feature thought to explain the low level of neutral genetic differentiation among populations



Fig. 1 Expected effects of random and directional gene flow between locally differentiated populations on individual genetic values. (a) In the case of random gene flow, migrant gene flow (orange arrows) tends to homogenize the frequencies of the red, grey and blue genotypes in each environment. Hence, in the core population (e.g. grey habitat), individuals and gametes from migrant gene flow should exhibit greater genetic variance for the traits under local selection ($v_1 > v_2$) than individuals and gametes from local gene flow (i.e. gene flow within population, represented by the green arrows), whereas the mean genetic value (dotted line) should be identical (assuming a gradual shift of the optimum). (b) In the case of directional gene flow (e.g. only the red genotypes migrate to the grey habitat), migrant gene flow should introduce individuals and gametes with shifted mean genetic values (dotted lines; $m_1 \neq m_2$), whereas the variance in genetic values between migrant and local gene flow should be similar (assuming that stabilizing selection is the same in each habitat).

New Phytologist (2020) www.newphytologist.com observed from the local to the European scale (Comps *et al.*, 2001; Buiteveld *et al.*, 2007). Several phenotypic traits show significant genetic differentiation among beech provenances at the European scale (Robson et al., 2018), in particular budburst date (Gomory & Paule, 2011; Kramer *et al.*, 2017), height (Rose *et al.*, 2009) and, to a lesser extent, wood anatomy, hydraulic traits and physiological foliar traits (Knutzen *et al.*, 2015; Hajek *et al.*, 2016; Stojnic *et al.*, 2018). These patterns are usually interpreted as a long-term response to selection, conferring better adaptation of beech populations to local climate and, most importantly, to summer drought. Our study aimed to test whether such patterns of phenotypic adaptation also exist at a fine spatial scale.

Here, we analyzed the phenotypic and genotypic data of \sim 2300 beech seedlings from 60 maternal families grown in a common garden. These open-pollinated families were collected from three natural beech plots, spreading along an elevation gradient at fine spatial scale (c. 1.5 km in-between the most distant plots) at the southern, warm margin of this species' distribution. We focused on 11 potentially adaptive phenotypic traits, for which we previously found a significant genetic component in phenotypic variation (Gauzere et al., 2016a). The first question investigated was: (1) Is there genetic divergence across elevations for the quantitative traits of interest? We expected trees from the low-elevation, warmer and drier plot to have evolved traits favouring adaptation to drought, and trees from the high-elevation, colder plot to have evolved traits favouring adaptation to low temperatures. Our second question was: (2) Is there a signature of local selection in the differentiation pattern of the trait studied. We tested this by comparing traits differentiation to neutral differentiation shown by microsatellite markers. Our final question was: (3) Does gene flow contribute to the additive variance of traits under selection at the local scale? To that purpose, we used parentage



Fig. 2 Climatic space explored by the three study plots in comparison to the whole species range of *Fagus sylvatica*. This climatic space is represented as the variation of temperature and summer precipitation over the beech distribution area, extracted from the SAFRAN database for the period 1958–2015 (collected on a 8 km² grid represented by grey dots). Each black triangle represents the average climate recorded since 2007 with HOBO weather stations for plots N1 (1020 m above sea level (asl)), N2 (1140 m asl) and N4 (1340 m asl).

analysis to identify the offspring likely originating from local or distant pollen immigration events, and we tested how long-distance pollen dispersal affected the genetic values of offspring after one episode of reproduction.

Materials and Methods

Species, sampling plots and experimental design

Fagus sylvatica L. (European beech) is a monoecious, anemophilous and predominantly outcrossed tree species (Merzeau et al., 1994; Wang, 2003). It is a shade-tolerant species requiring well-drained and moderately deep soils and relatively high humidity rates (Breda et al., 2006; Jump et al., 2006). Its distribution ranges from the northern Mediterranean regions to the south of Scandinavia. On Mont-Ventoux, a mountain in southeastern France, beech forests are located at the southern limit of their ecological range (Fig. 2), in a mountainous Mediterranean ecosystem. On the northern face of Mont-Ventoux, the beech forest ranges almost continuously from 750 to 1700 m above sea level. This steep elevation gradient provides almost linear variation in mean temperature and humidity with elevation (Davi et al., 2011). Three plots were defined along this climatic gradient, at 1020 m above sea level (asl) (N1), 1140 m asl (N2) and 1340 m asl (N4) (Supporting Information Table S1), over a total distance of c. 1.5 km. A previous study showed that these plots have high pollen immigration rates (on average 53.3%), moderate distances of pollen dispersal (on average 45.7 m), and non-negligible long-distance pollen dispersal events (Gauzere et al., 2013b).

In August 2009, 20 open-pollinated families were sampled in each of the three plots (60 maternal progenies in total). Mothertrees were chosen for their high fertility and random location within the plot. In April 2010, the seeds were germinated and a subset of 46.4 seedlings on average per family were randomly planted in 25 blocks in a common garden experiment (with 1.8 seedlings per family in each block) at the State nursery of Aix-en-Provence (lat. 43°23'30"N, long. 5°23'24"E). All seedlings were planted in independent pots of 1.2 l with sand substrate and fertilizer, and irrigated regularly. The experiment ran for 3 years (from April 2010 to September 2013). Details on the seed collection, germination protocol and progeny test design can be found in Gauzere *et al.* (2016a).

All of the potentially reproductive adults within the three plots, including the mother-trees, and an average of 23.9 off-spring per family were genotyped at 13 microsatellites markers (690 adults and 1437 offspring in total; see Gauzere *et al.*, 2013b, for the genotyping details).

Phenotypic measurements

A set of 11 different phenotypic traits were measured on the offspring grown in the common garden. These traits, already introduced in Gauzere *et al.* (2016a), are presented briefly below (see also Table 1). Height and diameter growths were measured between August 2010 and November 2011 (ΔH , ΔD). For the

Table 1 Description and descriptive statistics for the traits measured on seedlings of Fagus sylvatica.	Table 1	Description and	descriptive statistics	for the traits measure	d on seedlings of	Fagus sylvatica.
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Abbreviation	Trait	Category	N _w	Unit	Transformation
ΔH_{\log}	Total height increase	Growth	2380	Rate	log
ΔD	Total diameter increase	Growth	2380	Rate	_
$t_{b2 \rightarrow 3(2011)}$	Budburst date in 2011	Phenology	2334	DOY	_
$t_{b2\to 3(2012)}$	Budburst date in 2012	Phenology	1835	DOY	_
$t_{s1\rightarrow 2}$	Senescence date in 2011	Phenology	2386	DOY	_
VD ₂₀₁₁	Vegetation season duration in 2011	Phenology	2193	DOY	_
LM	Dry leaf mass	Morphology	2491	g	Square-root
LA	Leaf area	Morphology	2491	cm ²	Square-root
LMĂ	Leaf mass area	Morphology	2491	$\mathrm{g}\mathrm{m}^{-2}$	
% N	Nitrogen content in leaves	Physiology	1031	%	_
$\delta^{13}C$	Water use efficiency	Physiology	1031		_

DOY, day of the yr. N_{w} , number of phenotyped individuals.

whole trial, phenological events also were recorded: the date of budburst in 2011 and 2012 ($t_{b2\to3(2011)}, t_{b2\to3(2012)}$), the date of leaf senescence in 2011 $(t_{s1\rightarrow 2(2011)})$ and the duration of the growing season in 2011 (VD₂₀₁₁ = $t_{s1 \rightarrow 2(2011)} - t_{b2 \rightarrow 3(2011)}$). At the end of the growing season in 2011, three light-exposed leaves were collected on the stem of each seedling to measure morphological and physiological traits. On fresh leaves, we first measured the leaf area (LA), and put them to dry at 60° C to then measure the leaf dry mass (LA) of each seedling. The leaf mass area, related to the photosynthetic capacity and stomatal conductance of the plant (Reich *et al.*, 2003), was calculated as LMA = $\frac{LM}{LA}$. Finally, for a subset of 1031 individuals (evenly sampled across families), we measured the leaf carbon isotope composition $(\delta^{13}C)$, as a surrogate for water use efficiency (Farquhar *et al.*, 1989). The nitrogen content in the leaves (%N) also was estimated for this subset of individuals, as leaf nitrogen density is linked to the photosynthetic capacity of leaves (Kattge et al., 2009). Note that three of these traits were transformed before analyses to limit departure from linear model assumptions: ΔH with a logarithm transformation, and LM and LA with a squareroot transformation (following Gauzere et al., 2016a).

Test for population genetic differentiation

In order to assess the genetic differentiation of quantitative traits among plots we ran a specific linear mixed model (called 'animal model') that uses relatedness information from a pedigree to dissociate the contribution of genetic, population and environmental effects on the total phenotypic variance of traits. Thus, we analyzed each trait Y measured in common garden using the following model, and focused particularly on the significance of the fixed plot effect P:

$$Y_{i,f,p,k} = \mu + P_i + B_f + M_p + a_k + \varepsilon_{i,f,p,k}$$
Eqn 1

with, as fixed effects, μ the intercept, P_i the effect of the plot of origin, B_f the effect of the block and M_p the observer effect (only included for the analysis of the phenological traits). The random terms are the additive genetic values of the individuals k,

 $\{a_k\}: N(0, AV_A)$ with A the matrix of pairwise relatedness derived from a pedigree, and V_A is the additive genetic variance, and the residuals, $\{\varepsilon_{i,f,p,k}\}: N(0, I_{de}V_R)$ with I_{de} the identity matrix and V_R the residual variance.

The (co)variance structure of the additive genetic effects was defined using a one-generation pedigree reconstructed in a previous study (Gauzere *et al.*, 2016a). Here, we did not model maternal effects because of the lack of power to accurately dissociate maternal and additive genetic effects (Gauzere *et al.*, 2016b). Model Eqn 1 was fitted in R/ASREML (Gilmour *et al.*, 2006). Using a similar model, Gauzere *et al.* (2016a) previously showed that all traits presented moderate narrow-sense heritability, with h^2 ranging from 0.12 to 0.35.

Signature of natural selection accounting for genetic drift

We used the method developed by Ovaskainen *et al.* (2011) to detect signatures of natural selection in trait differentiation among populations. This method requires genotypic and phenotypic data, that are first analysed under a neutral model which assumes that the genes coding for the trait are as divergent as the neutral markers ($Q_{st} = F_{st}$ case). This first model thus considers the population mean genetic additive values as normally distributed, with the covariance between pairs of population means being proportional to the pairwise average coancestry between these populations (i.e. assuming neutrality for the trait):

$$\mathbf{A}^{\mathbf{P}}: N(0, 2V_{\mathbf{A}} \times \boldsymbol{\theta}^{\mathbf{P}})$$
 Eqn 2

with $\mathbf{A}^{\mathbf{P}}$ the vector of the population-level effects, V_{A} the additive genetic variance of the trait, $\mathbf{\theta}^{\mathbf{P}}$ the matrix of pairwise population-level coancestry coefficients. Note that the $\mathbf{\theta}^{\mathbf{P}}$ matrix provides an estimation of F_{st} .

The method then considers an animal model for each quantitative trait Y, that decomposes the additive value into a population and an individual effect. The population-level additive effect is determined by the mean genetic additive values in the parent population and the individual-level additive effects a_k are defined the same way as in model Eqn 1. The presence of selection on the trait is tested with the *S* statistics, which evaluate whether the realized pattern of population mean genetic additive values is likely under the neutral model Eqn 2.

In order to apply this method to our dataset, we first used the genotyping of the adult trees and the admixture *F*-model for neutral divergence among populations implemented in R/RAFM, to estimate θ^{P} and the index of neutral genetic differentiation, F_{st} (Karhunen & Ovaskainen, 2012). We then analyzed the traits measured in the common garden using the same fixed effects as in model Eqn 1 and the one-generation pedigree using R/DRIFT-SEL, to estimate V_A , \mathbf{A}^P and *S* (Karhunen *et al.*, 2013). These Bayesian analyses were repeated three times per trait to ensure convergence of the Markov chain Monte Carlo (MCMC) chains. We provide the average posterior *S* estimates (the chains provided very similar *S* values; Table S2). We also measured a low autocorrelation within the chains, indicating that the space of the parameter values is explored efficiently.

Immigrant characterization based on molecular markers

Categorical assignments of paternity We first tried to identify the father of the genotyped seedlings to characterize their immigration status. To that purpose, we used the likelihood-based method of categorical paternity assignment implemented in CERVUS (Marshall et al., 1998) with allelic frequencies estimated from the genotypes of all adult trees. All of the reproductive adults from the three sampled plots were considered as candidate fathers, but only fathers within the mother's plot were assigned (see below). We considered 0% typing error, 100% sampling of candidate fathers, a confidence level of 95%, and allowed selfing (following Gauzere et al., 2016a). These choices were made to favour assignments to genotyped fathers, despite the risk of type I errors (i.e. a wrong tree is assigned while the true father is not sampled), knowing that quantitative genetic methods are robust to pedigree errors (Charmantier & Réale, 2005; Gauzere et al., 2016b). We successfully assigned the paternity for 45.8% of the genotyped offspring. We considered that individuals assigned to a father located within their mother's plot come from 'local' pollen pool, whereas individuals with no compatible father were considered 'immigrant'. Noncategorized individuals include those genotyped at fewer than six markers, offspring not assigned because of compatibility with multiple fathers, and offspring assigned to a father outside their mother's plot. Indeed, given the large number of trees potentially contributing to median- and long-distance mating events, we considered that our ability to retrieve the true father outside the mother's plot was low and presented large expected type I error rate.

Probability of long-distance pollination In order to refine the 'immigrant' status, we also used the whole genetic dataset and a noncategorical paternity analysis to assess the likelihood for each genotyped seedlings to belong to long-distance immigrant pollen pool. To that purpose, we estimated the Mendelian transition probabilities (*T*) that each offspring has the observed genotype g_0 while originating from cross-fertilization of female j with an ungenotyped male from a population with the allelic frequencies

BAF: $T(g_0|g_j, BAF)$. We estimated this transition probability using (1) the global allelic frequencies estimated from a largerscale study on the whole north-face of Mont-Ventoux (AF_{global}; Lander *et al.*, 2011), and (2) the local allelic frequencies estimated from the genotyping of all the reproductive trees in the three study plots (AF_{local}). This allowed us to compute the probabilities that each offspring has the observed genotype g_0 given that it originates from long-distance and local pollination events, $T(g_0|g_j, AF_{global})$ and $T(g_0|g_j, AF_{local})$, respectively. From these two probabilities, we derived for each offspring the logarithm of the odds (LOD)-score of long-distance immigration vs local immigration:

$$v_{g_0} = \log\left(\frac{T(g_0|g_j, AF_{global})}{T(g_0|g_j, AF_{local})}\right)$$
Eqn 3

 $v_{g_0} > 0$ indicating that offspring g_0 is more likely originating from a long-distance pollination event and $v_{g_0} < 0$ that it is more likely originating from a short-distance pollination event.

These transitions probabilities were estimated using the MEMMi model (Gauzere *et al.*, 2013b).

Impact of pollen immigration on quantitative traits

We explicitly tested whether (1) offspring originating from distant fathers presented more genetic variance for quantitative traits than offspring originating from local fathers, which is expected in the case of migration from genetically differentiated populations in an island model, and whether (2) offspring originating from distant fathers had shifted average genetic values for quantitative traits as compared to the ones originating from local fathers, which is expected in the case of directional pollen immigration from genetically differentiated populations along the gradient (Fig. 1). In both cases (1) and (2), pollen dispersal would result in increasing genetic variance and adaptive capacities. We ran two different models depending on how the immigration status was characterized (categorical or quantitative variable).

First, for each phenotypic trait *Y*, we looked for differences in the mean and variance of additive genetic values (a_k) between 'local' and 'immigrant' groups, using the following animal model:

$$Y_{i,f,p,k} = \mu + P_i + B_f + M_p + a_k + \varepsilon_{i,f,p,k}$$
 Eqn 4

with the same fixed effects than in model Eqn 1, but here the (co)variance structure of the additive genetic effects depend on the immigration classes, with l= 'local' and m= 'immigrant':

 $\begin{bmatrix} V_{Al} & 0\\ 0 & V_{Am} \end{bmatrix}$. Note that the effect of 'local' and 'immigrant' cat-

egories on the average trait values was tested and found to be never significant. This fixed effect is thus absent in the final model.

From model Eqn 4, we thus estimated two variance components: $V_{\rm Al}$ and $V_{\rm Am}$.

We then performed a random regression model to investigate the effect of long-distance pollen dispersal, *v*, on the variation in additive genetic variance. Random regression models are commonly used in evolutionary biology and breeding to allow additive genetic effects (or any random effect) to vary with a covariate (e.g. environment, age; Nussey *et al.*, 2008). This model defines a random intercept and slope at the additive genetic level, as deviation from the mean linear effect of v on Y, to model the variation of V_A with v as: $V_A(v) = V_{Aintercept} + v^2$. $V_{Aslope} + 2 \cdot v \cdot cov_A$ (intercept, slope). The effects of v on V_A was tested using the following models:

$$Y_{i,f,p,k} = \mu + P_i + B_f + M_p + \alpha v_k + a_k + \varepsilon_{i,f,p,k}$$
 Eqn 5

$$Y_{i,f,p,k} = \mu + P_i + B_f + M_p + \alpha v_k + \tilde{a}_k + \varepsilon_{i,f,p,k}$$
 Eqn 6

where $\tilde{a}_k = a_k + \beta_k v_k$ describes the random intercept and slope of v on Y at the additive genetic level in model Eqn 6. In both models Eqn 5 and 6 the term αv_k is the fixed effect of the LODscore of long-distance vs local migration on the trait.

From the model Eqn 6, we estimated the variances of the random intercept and slope, and the covariance between them. Note that for each individual k, we only have one value of v and Y, but the model estimates these (co)variance components by comparing phenotypic similarities and the effects of v on Y between relatives.

The significance of the qualitative and quantitative effects of pollen immigration on V_A were tested by comparing the two hierarchical models Eqn 1 and 4, and the two models Eqn 5 and 6, with a log-likelihood ratio test. We performed these analyses with R/ASREML, considering only the information about maternal relatednesses, as we (often) ignore the father's identity of 'immigrant' individuals.

Relatedness and Nep estimates

We used the genetic dataset to estimate the average paternal relatedness ($\overline{\rho}_p$) and effective number of pollen donors ($Nep = 1/\overline{\rho}_p$) in the 'local' and 'immigrant' pollen clouds. We first extracted the haplotype corresponding to the paternal contribution to each diploid genotype following Gauzere *et al.* (2016a). These paternal genotypes were then used to estimate the pairwise paternal relatednesses among each pair of offspring *k* and k' ($\rho_{k,k'}$) using the kinship coefficient of Loiselle *et al.* (1995) implemented in SPAGEDI (Hardy & Vekemans, 2002), and the local allelic frequencies (AF_{local}). These coefficients were estimated removing all the selfed individuals to only compare the outcrossed pollen pools.

Results

Neutral genetic differentiation among plots and pollen immigration

The admixture *F*-model of Karhunen & Ovaskainen (2012) estimated a posterior F_{st} value of 0.026 [0.024; 0.029], consistent with previous estimates of neutral genetic differentiation among populations measured with a comparable method but from 51 populations over a much wider area of Mont-Ventoux ($F_{st} = 0.026$; Lander *et al.*, 2011). The coancestry estimates within plots (diagonal elements of the $\theta^{\rm p}$ matrix) indicated that plot N1 tended to have lower effective population size than plot N2 and N4 (Table 2). Plots N1 and N2 exhibited slightly higher between-plots coancestry coefficient, suggesting that the two closest plots exchanged more gene flow ($\theta_{\rm N1N2} > \theta_{\rm N1N4} \sim \theta_{\rm N2N4}$). Overall, the coancestry estimates were low (maximum $\theta_{\rm N1} = 0.057$; Table 2) indicating that effective population sizes were globally large. This result is consistent with the large effective number of pollen donors *Nep* estimated for these sites using the same dataset (Table S1; Gauzere *et al.*, 2013b).

Using paternity assignment, we categorized n = 658 offspring as 'local' (45.5%) and n = 492 as 'immigrant' (34%). A total of 295 (20%) offspring were outside of these two categories (i.e. ambiguous origin). Note that this 'immigrant' status includes offspring with nonsampled fathers which could be located either at a short or long distance to the mother-tree's plot. We checked that the LOD-score of long-distance vs local immigration was indeed higher on average for the 'immigrant' than 'local' individuals, with $\overline{v} = -0.33$ and 0.18 for local and immigrant offspring, respectively (*P*-value < 0.001, ANOVA test; Fig. S1).

The analysis of the pairwise coefficient of paternal relatedness showed that within the immigrant pollen pool fathers were significantly less related than within the local pollen pool, with $\overline{\rho}_{\rm p} = 0.011$ and $\overline{\rho}_{\rm p} = 0.007$ within the local and immigrant pollen pools, respectively (*P* < 0.001, ANOVA). Consequently, we estimated a higher effective number of pollen donors (*Nep*) in the immigrant than in the local pollen pool (*Nep* = 26.4 and 31.3 for local and immigrant pollen pools, respectively).

Patterns of genetic trait differentiation and test for selection

Significant genetic differentiation across elevations was found for ΔH_{\log} , ΔD , $t_{b2\rightarrow3(2011)}$ and $t_{b2\rightarrow3(2012)}$, whereas marginal differentiation was found for $t_{s1\rightarrow2(2011)}$ (as revealed by the *P* effect in model Eqn 1; Fig. 3). Using the method of Karhunen & Ovaskainen (2012), we detected a signal of natural selection for three of these four traits: ΔH_{\log} , ΔD and $t_{b2\rightarrow3(2011)}$ (i.e. S > 0.90; Fig. 3). For growth traits, plots at intermediate and high elevations presented the highest height and diameter increments

Table 2 Matrix of the coancestry coefficients estimated by the admixture*F*-model of Karhunen & Ovaskainen (2012) for the three plots of *Fagussylvatica*.

	N1	N2	N4
N1	0.057 [0.033; 0.085]		
N2	0.018 [0.005; 0.035]	0.029 [0.017; 0.045]	
N4	0.012 [0.00; 0.024]	0.013 [0.002; 0.023]	0.035 [0.02; 0.052]

The diagonals correspond to the within-plot coancestry coefficient, with its median value and confidence intervals (i.e. values of the estimates that bound 95% of the posterior distribution). Lower diagonal elements are the median coancestry coefficients among-plots, with confidence intervals in brackets.

 $(\Delta H_{\log N2} = 0.88, \Delta H_{\log N4} = 1.03 \text{ and } \Delta D_{N2} = 0.82, \Delta D_{N4} = 0.72)$, whereas the site at the lowest elevation showed the lowest growth $(\Delta H_{\log N1} = 0.56 \text{ and } \Delta D_{N1} = 0.50; \text{ Fig. 3})$. For the phenological traits, buds flushed earlier for plots from high elevations as compared to low elevation plots $(t_{b2\rightarrow3(2011)N1} = 101.9; t_{b2\rightarrow3(2011)N2} = 100.3; t_{b2\rightarrow3(2011)N4} = 99.8; \text{ Fig. 3})$. Interestingly, contrary to growth and phenological traits, morphological and physiological traits (except LMA) tended to be slightly less differentiated than expected under neutral processes (with $S \in [0.24; 0.39]$), with a strong conservation of the genetic values between plots (e.g. for the leaf mass $LM_{\sqrt{N1}} = 0.44$, $LM_{\sqrt{N2}} = 0.45$, $LM_{\sqrt{N4}} = 0.44$; Fig. 3).

Effect of pollen immigration on within-plot genetic variation

Using the categorical assignment and the model Eqn 4, we only estimated marginally significant difference in V_A between 'local' and 'immigrant' individuals for $t_{b2\rightarrow3(2012)}$ (P=0.088; for all of the other traits P > 0.1). In that case, we predicted higher V_A within the 'immigrant' than the 'local' group ($V_{A_{immig}} = 16.57$ and $V_{A_{local}} = 3.16$; Table 3). Using the random regression model Eqn 6, we found significant variation in additive genetic variance with the likelihood of provenance from long-distance pollen pool for ΔH_{\log} (P=0.005), and a marginally significant effect for $t_{b2\rightarrow3(2012)}$ (P=0.078; for all the other traits P>0.1; Table 3). The estimated variance-covariance terms showed increasing $V_{\rm A}$ with higher likelihood of long-distance pollination event for both ΔH_{\log} and $t_{b2 \rightarrow 3(2012)}$ (Fig. 4). Nevertheless, for these traits, the fixed effect of v was not significant, which indicated no global trend in the variation of breeding values with the likelihood that an offspring originates from long-distance pollen dispersal. Interestingly, these two different analyses highlighted the same trend towards a higher genetic variance for offspring originating from long-distance dispersal than from local dispersal.

Discussion

The scale of phenotypic adaptation

This study provides evidence that divergent selection can act on phenotypic traits at fine spatial scale in temperate tree species. The study beech plots showed important dispersal capacities, with pollen dispersal that can easily reach 100 m, a fat-tailed pollen dispersal kernel and high rates of immigration (Gauzere *et al.*, 2013b; Oddou-Muratorio *et al.*, 2018), similar to other anemophilous tree species (see Ashley, 2010; Kremer *et al.*, 2012, for reviews). Because dispersal distances overlap the spatial scale of the selective environment, our results provide strong evidence that adaptive genetic differentiation can occur despite high gene flow (see Fitzpatrick *et al.*, 2015; Peterson *et al.*, 2016, for recent evidence in other taxa). This also suggests that selective pressures on seedling growth (in diameter and height) and on budburst timing were strong enough to counteract the homogenizing effect of gene flow on structuring

the genetic variation at the scale of a few hundred metres up to one kilometre. We found that the seedlings derived from the plots at high elevation, inhabiting the lowest temperature conditions, flushed earlier and had a higher height and diameter growth in the common garden than the plot at low elevation. For the budburst date, this pattern of genetic differentiation is opposed to the pattern of phenotypic variation measured in situ, where seedlings from plots at low elevations flushed earlier than plots at high elevations (Davi et al., 2011). Beech populations originating from higher longitude or elevation also have been shown to be genetically earlier in provenance tests (Teissier du Cros et al., 1988; Gomory & Paule, 2011; Kramer et al., 2017), suggesting that these populations evolved phenological traits promoting a longer growing season length. In forestry, growth is considered a good approximation of the performance of trees in a given condition (e.g. Rehfeldt et al., 1999, 2002). In situ and ex situ, the highest diameter growth values were measured at intermediate to high elevations (Cailleret & Davi, 2011, and present study), suggesting that the genetic and environmental influences on growth traits are in the same direction. At this southern edge of the ecological conditions tolerated by beech (Fig. 2), the reduced allocation to stem growth at the low-elevation plot is likely an adaptive response to drought, which has been described previously by comparing marginal vs central beech populations (Rose et al., 2009). These studies, including ours, shed light on the phenotypic traits under selection and underlying adaptation at different spatial scales. However, they do not provide a direct assessment of the strength of natural selection or the ecological bases of selection, nor do they quantify phenotypic mismatch and maladaptation, which would require selection analyses (MacColl, 2011).

The weak adaptive divergence of morphological and physiological traits as compared to growth and phenological traits along ecological gradients remains an open question. Meta-analyses highlighted that growth and phenological traits are the most genetically differentiated phenotypic traits at large spatial scale in temperate and boreal tree species (Savolainen et al., 2007; Alberto et al., 2013). Here, our study suggests that this conclusion also holds at a narrower spatial scale. The existing literature also tend to indicate that morphological and physiological traits generally exhibit a lower fraction of genetically controlled phenotypic differentiation, and a higher within-population genetic variation than other traits (Hajek et al., 2016). A putative explanation is that latitudinal or elevation gradients do not fully capture the spatial scale of variation for drought stress, even if they capture a strong variation in temperature. In particular in our study, the important microheterogeneity in soil water capacity on (Nourtier et al., 2013) may generate micro-local differentiation of morphological and physiological traits, blurring the signal of divergent selection for morphological and physiological traits at a larger spatial scale (Cubry et al., unpublished). Another possible explanation is that morphological and physiological traits could be under stabilizing selection (e.g. Lamy et al., 2011). Unfortunately, the power to detect the signal of stabilizing selection is low for species with low neutral genetic differentiation, such as

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Fig. 3 Level of trait adaptive divergence between the three plots located at 1020 m above sea level (asl) (N1), 1140 m asl (N2) and 1340 m asl (N4) for the growth, phenological, physiological and morphological traits measured on seedlings of *Fagus sylvatica*. Histograms represent the *a posteriori* distributions of additive genetic values estimated for each plot. The dotted lines represent the median of these distributions. We also provide the test for population differentiation (*P*) and signal of selection (*S*) for these quantitative traits. An *S* coefficient close to 1 and 0 indicates signal of divergent and stabilizing selection, respectively. ***, P < 0.001; **, 0.001 < P < 0.05; ., 0.05 << P < 0.09; ns, P > 0.09.

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Table 3 Effect of pollen immigration on the variation in additive genetic
variance for traits measured in seedlings of Fagus sylvatica.

Model Eqn 4		
Trait	V _{Aimmig}	V _{Alocal}
$t_{b2 \rightarrow 3(2012)}$	16.57 [ँ] (8.2)	3.16 (5.6)
Model Eqn 6		
Additive genetic effects for	or ΔH_{\log}	
	$\Delta H_{\log_{intercept}}$	$\Delta H_{\log_{slope}}$
$\Delta H_{\log_{intercept}}$	0.16 (0.041)	osope
$\Delta H_{\log_{slope}}$	0.029 (0.010)	0.0092 (0.008)
Additive genetic effects for	or $t_{b2 \to 3(2012)}$	
	$t_{b2 \rightarrow 3(2012)_{intercept}}$	$t_{b2 ightarrow 3(2012)_{slope}}$
$t_{b2 \rightarrow 3(2012)_{intercept}}$	7.36 (4.8)	
$t_{b2 \rightarrow 3(2012)_{slope}}$	3.44 (2.12)	2.47 (2.18)

This effect was found significant or marginally significant for two genetically differentiated traits: using model Eqn 4 for $t_{b2\rightarrow3(2012)}$ with P = 0.088; using model Eqn 6 for ΔH_{log} with P = 0.005 and for $t_{b2\rightarrow3(2012)}$ with P = 0.078. Model Eqn 4 estimates an additive genetic variance (V_A) for each 'immigrant' and 'local' group (SE in brackets). Model Eqn 6 estimates the additive genetic variance-covariance terms for the intercept and slope of the trait with the logarithm of the odds (LOD)-score of long-distance immigration vs local immigration v (SE in brackets). The diagonal elements thus provide V_{Aintercept} and V_{Aslope}, and the off-diagonal element cov_A (intercept, slope). See Supporting Information Tables S3 and S4 for the model outputs for all phenotypic traits.

trees (Savolainen *et al.*, 2007), and patterns of weak quantitative trait differentiation are difficult to interpret (Whitlock, 2008). With the development of genomic resources, this question of microgeographic adaptation is now addressed mainly using genome-wide association studies (e.g. Eckert *et al.*, 2015). However, associations between genotype, phenotype and environment often provide weaker evidence of adaptive differentiation at the phenotypic levels (Le Corre & Kremer, 2012), probably because of the complex multilocus determinism of the traits underlying phenotypic adaptation. Using a more precise characterization of ecological gradients (e.g. by considering the spatial-scale of water

availability) and new methods to compare the similarity of habitats and phenotypes offer promising prospects to better understand the spatial scale of phenotypic adaptation and the ecological drivers of adaptive divergence (Karhunen *et al.*, 2014; Csillery *et al.*, 2020).

Effects of pollen dispersal on quantitative trait variation at a local scale

For populations locally adapted and experiencing strong selective pressures that vary through space, theory suggests that moderate gene flow can maintain quantitative genetic variation within these populations (Barton, 2001; Le Corre & Kremer, 2003; Bridle et al., 2010). Although adaptive genetic differentiation is widespread, only a few studies have provided evidence that gene flow is an important process to maintain the high levels of genetic variation usually detected in nature (e.g. Yeaman & Jarvis, 2006). Here, consistent with this theoretical expectation, we detected an effect of pollen dispersal on increased genetic variance for two locally differentiated traits: height growth and budburst date. However, we likely lacked the power to detect this effect for the third heritable trait under selection - diameter growth. Because female flowers are receptive before the emission of local pollen (protogyny) and lower populations flower earlier than higher populations, we hypothesized a directional pollen flow across elevations in Gauzere et al. (2013a), which would have resulted in shifted genetic values between immigrant and local individuals (Fig. 1; Nilsson, 1995). Our results did not support this expectation.

We used two different approaches to characterize the pollen immigration status. Both analyses suggested that immigrant pollen brings more genetic variance for locally differentiated traits than local pollen. These effects were quantitatively high, with genetic variance for immigrants more than twice as high as for local offspring. However, this result had large standard errors



Fig. 4 Change in additive genetic variance (VA) with the logarithm of the odds (LOD)-score of long-distance immigration vs local immigration (v), estimated using model Eqn 6 in *Fagus sylvatica*. This relationship is represented only for two of 11 traits tested, for which we found that this effect was significant or marginally significant: (left) the log-transformed height growth (ΔH_{log}) and (right) the budburst date in 2012 ($t_{b2\rightarrow3(2012)}$). The solid line represents the estimate, and the dashed lines the 95% confidence intervals around this estimate. From the random regression model Eqn 6, we can estimate the change in V_A with a covariate x as: $V_A(x) = var(trait_{intercept} + trait_{slope} \cdot x) = V_{Aintercept} + x^2 \cdot V_{Aslope} + 2 \cdot x \cdot cov_A(intercept, slope)$, using the estimates provided in Table 3.

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around the estimates, and the significance changed depending on how the immigration status was characterized. We checked that 'immigrant' individuals originated from more diverse fathers (higher effective number of pollen donors Nep), and had paternal genotypes more similar to global than local allelic frequencies (higher LOD-score of long-distance vs local immigration, v). Nevertheless, our immigrant status still likely suffers some inaccuracy. In particular, the low neutral genetic differentiation among the study beech plots ($F_{st} = 0.026$) and, overall, on the northern face of Mont-Ventoux (Lander et al., 2011), may reduce our ability to retrieve the origin of a pollen grain using molecular markers. Moreover, the low levels of quantitative genetic differentiation measured along the elevation gradient suggest that the expected effect of pollen flow on genetic values after one episode of reproduction is necessarily moderate. Here, this effect may thus be difficult to detect, even knowing the true immigration status of all offspring.

The approach used here provides a direct quantification of the impact of contemporary pollen immigration on the genetic variance of quantitative traits. Although using molecular markers to distinguish migrants and local offspring could have been done on numerous existing datasets, to our knowledge no study before ours had reiterated the approach first used by Nilsson (1995). However, we cannot estimate the immigration load associated with this increase in genetic variance for the locally differentiated traits. Although this genetic load is expected to be beneficial under rapidly changing environmental conditions (Yeaman & Jarvis, 2006; Savolainen et al., 2007; Kremer et al., 2012), this remains to be tested. Applying this approach to other species and populations may improve our ability to both understand and quantify the impact of gene flow on the adaptive potential of quantitative traits under natural conditions. This effect should be more detectable for populations departing from migration-selection equilibrium, such as isolated (or marginal) populations, or transplanted plant populations (see Fitzpatrick et al., 2015, 2016, for examples in guppies).

The present results about trait differentiation and genetic variance of migrants and nonmigrants extend previous studies about pollen dispersal kernels and migration rates (Gauzere et al., 2013a), trait heritabilities (Gauzere et al., 2016a) and variance of fecundities (Oddou-Muratorio et al., 2018). Pulling all effects together suggests building a predictive mechanistic model to both (1) check whether the observed levels of genetic variance in migrant pollen are consistent with the amounts of migrant genetic variance that would be predicted mechanistically and (2) further investigate the adaptive dynamics at the scale of few generations (following, e.g., Kuparinen et al., 2010). This work is beyond the scope of the present study because it still requires a map of beech density over the whole northern face of Mont-Ventoux and an extrapolation model to propose a spatially explicit distribution of adaptive diversity. Point (2) would require information about the variation in selection gradients or adaptive landscapes with elevation. Unfortunately, empirical estimates of selection need specific approaches in natural populations of longlived species and, hence, remain scarce in forest trees (but see Bontemps et al., 2017).

Gene flow in the context of climate change

One of the most documented consequences of climate change on tree ecosystem functioning is probably the impact of climate warming on the advancement of spring phenology (Menzel et al., 2006; Fu et al., 2015). Understanding the adaptive or maladaptive value of the response of phenology to climate has become a key focus of evolutionary studies (Tansey et al., 2017; Gauzere et al., 2020). An underrated issue is that these phenological changes also are likely to affect gene flow among populations, and the efficiency of long-distance pollen dispersal. For many temperate tree species, we now have strong evidence that climate warming reduces spring phenology synchrony among individuals experiencing the same environmental conditions (Zohner et al., 2018), but also leads to more uniform spring phenology across climatic gradients (Vitasse et al., 2018). The evolutionary consequences of pollen dispersal may therefore change with increasing climate warming. In this study, we detected an effect of local selection and pollen dispersal on the adaptive potential of traits that are known to affect dispersal or reproduction in anemophilous tree species (tree height and phenology; Klinkhamer et al., 1997; Gauzere et al., 2013a; Oddou-Muratorio et al., 2018). In particular, long-distance pollen dispersal events are only effective if the donor and recipient trees have synchronous phenology (Aitken et al., 2008). Our results thus suggest that refining the models of phenotypic adaptation by incorporating the feedbacks between traits under divergent selection and dispersal-reproductive capacities may be important to thoroughly understand the evolutionary consequences of gene flow in plant species (e.g. Soularue & Kremer, 2012).

With the concern that populations could be maladaptated and lag behind with increasing climate change, new conservation and management strategies are needed to mitigate these effects. Assisted gene flow - the managed movement of individuals or gametes between populations within species ranges - is one of the tools proposed to help populations adapt to new climatic conditions (Aitken & Whitlock, 2013; Aitken & Bemmels, 2016). However, assisted gene flow among distant populations, at different latitudes, may have unexpected consequences for the response of individuals or new hybrids, because of outbreeding depression (Aitken & Whitlock, 2013), but also because environmental cues other than temperature can govern physiological responses in trees (see, for instance, the role of photoperiod in bud development; Way & Montgomery, 2015). Here, our results suggest some effect of local pollen dispersal on increasing genetic variation for locally differentiated traits after solely one episode of reproduction. Therefore, programmes considering the movements of gametes along short ecological gradients could be efficient to boost genetic diversity and introduce genetic innovation within natural or managed tree populations. Our results also encourage the development of evolution-orientated forest management and silviculture, whereby thinning could be done to favour the spread of long-distance gene flow (e.g. by reducing local densities; Lefevre et al., 2014). Nonetheless, unlike assisted gene flow, natural gene flow also can lead to increased maladaptation through migration load (Aitken & Whitlock, 2013).

Conclusion

By combining population genetics and quantitative genetics, we provided a comprehensive analysis of the effects of gene flow on the adaptive potential of a tree species in nature. We found that gene flow among trees at different elevations did not prevent adaptive differentiation among plots separated by < 1 km, and that divergent selection at this fine spatial scale acted mainly on growth and phenological traits, probably as an adaptation to drought and temperature variation. For two genetically differentiated traits, our analyses suggested an effect of pollen dispersal on increasing genetic diversity after one episode of reproduction. However, our power to detect and estimate these effects probably was quite low. Our results thus suggest that local selection is an important evolutionary force in natural tree populations and that, under rapidly changing environmental conditions, gene flow can increase the adaptive capacities of natural populations.

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Author contributions

JG, EKK and SO-M conceived and designed the study; JG collected the genetic and phenotypic data, performed the analyses, wrote the first draft of the text and contributed the revisions; SO-M conceived and coordinated the beech experiment; JG, HD, OB, EKK and SO-M discussed and interpreted the findings; and EKK, HD, OB and SO-M contributed to the analyses and the revisions of preliminary drafts.

Data availability statement

Data available upon request from the authors.

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Supporting Information

Additional Supporting Information may be found online in the Supporting Information section at the end of the article.

Fig. S1 Distribution of the LOD-score of long-distance vs local immigration between 'immigrant' and 'local' groups.

 Table S1 Location, climate and ecological characteristics of the study plots.

Table S2 Test for plot differentiation and signal of selection inquantitative traits.

Table S3 Differences in additive genetic variance between 'immigrant' and 'local' groups estimated from model Eqn 4.

Table S4 Effect of pollen immigration on the variation in addi-tive genetic variance estimated from model Eqn 6.

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